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        ANSWER 1 OF 1 CAPLUS COPYRIGHT 2006 ACS on STN
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        2004:182659
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        140:229441
TI
        Methods of reducing ischemic injury with compounds reducing activity of
        MAP kinase-activated protein kinase 2
IN
        Wang, Xinkang; Schieven, Gary; Feuerstein, Giora Z.
PA
        Bristol-Myers Squibb Company, USA
        PCT Int. Appl., 39 pp.
SO
        CODEN: PIXXD2
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L8
        ANSWER 1 OF 6
                                    MEDLINE on STN
ΤI
        Phorbol ester-induced G1 arrest in BALB/MK-2 mouse
        keratinocytes is mediated by delta and eta isoforms of protein kinase C.
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        ANSWER 2 OF 6 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights
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TI Phorbol ester-induced G1 arrest in BALB/MK-2 mouse keratinocytes is mediated by δ and η isoforms of protein kinase C.

- L8 ANSWER 3 OF 6 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on STN
- TI Phorbol ester-induced G1 arrest in BALB/MK-2 mouse keratinocytes is mediated by delta and eta isoforms of protein kinase C.
- L8 ANSWER 4 OF 6 SCISEARCH COPYRIGHT (c) 2006 The Thomson Corporation on STN
- TI Phorbol ester-induced G1 arrest in BALB/MK-2 mouse keratinocytes is mediated by delta and eta isoforms of protein kinase C
- L8 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN
- TI Phorbol ester-induced G1 arrest in BALB/MK-2 mouse keratinocytes is mediated by δ and η isoforms of protein kinase C
- L8 ANSWER 6 OF 6 BIOTECHNO COPYRIGHT 2006 Elsevier Science B.V. on STN
- Phorbol ester-induced G1 arrest in BALB/MK-2 mouse keratinocytes is mediated by δ and η isoforms of protein kinase C
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- L9 13 L1 AND ISCHEMIA
- => dup rem 19 PROCESSING COMPLETED FOR L9
- L10 8 DUP REM L9 (5 DUPLICATES REMOVED)
- => d ti 1-8
- L10 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- TI DEF domain-containing members of the MAP kinase pathway and their use in screening for drug inhibitors
- L10 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- TI Role of F-actin organization in p38 MAP kinase-mediated apoptosis and necrosis in neonatal rat cardiomyocytes subjected to simulated ischemia and reoxygenation
- L10 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- TI Methods of reducing ischemic injury with compounds reducing activity of MAP kinase-activated protein kinase 2
- L10 ANSWER 4 OF 8 MEDLINE on STN DUPLICATE 1
- TI Mitogen-activated protein kinase-activated protein (MAPKAP) kinase 2 deficiency protects brain from ischemic injury in mice.
- L10 ANSWER 5 OF 8 SCISEARCH COPYRIGHT (c) 2006 The Thomson Corporation on STN
- TI Stimulation of multiple MAPK pathways by mechanical overload in the perfused amphibian heart
- L10 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- TI Ischemic preconditioning triggers tyrosine kinase signaling: a potential role for MAPKAP kinase 2
- L10 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- TI Ischemic preconditioning: role of multiple kinases in signal amplification and modulation
- L10 ANSWER 8 OF 8 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on STN
- TI Involvement of a tyrosine kinase-dependent signal transduction process involving p-38 MAP kinases and MAPKAP kinase 2 in ischemic

preconditioning.

=> d ab 1 3 4 6 8 110

- L10 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- Mitogen-activated protein (MAP) kinases (e.g., ERK1/2) phosphorylate a variety of target proteins including, for example, several immediate-early gene products (e.g., Fos, Myc, and Jun family proteins). Certain phosphorylation reactions require binding of the MAP kinase to the DEF domain of the target protein. Inhibitors that block this interaction may be useful therapeutics for human disease, including as antineoplastic agents. This invention provides several advantages over known therapies that directly target the MAP kinase signaling cascade. Typically, most compds. that inhibit the MAP kinase pathway are non-specific and inhibit more than one enzyme, and the targeted inhibited kinases are not available to perform normal physiol. functions necessary for cell survival, whereas therapeutic methods of the present invention inhibit the activation of particular target proteins and leave the MAP kinases enzymically active and available to phosphorylate other non-DEF domain-containing proteins. Thus, DEF domains are identified in a large number of proteins, and the principles of the invention are exemplified using the immediate-early gene, c-Fos. Screening assays useful for identifying compds. that inhibit the MAP kinase-DEF domain interaction are also disclosed.
- L10 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- AB The present invention includes methods of reducing the activity, such as enzymic activity and expression, of mitogen-activated protein (MAP) kinase-activated protein kinase
 2 (MK2). The present invention further includes methods for identifying compds. useful for reducing such activity, and methods for reducing ischemic injury by the administration of such compds. Ischemic brain injury was significantly reduced in MK2 deficient mice compared to theat of wild type mice following either transient or permanent occlusion of the middle cerebral artery.
- L10 ANSWER 4 OF 8 MEDLINE on STN DUPLICATE 1
 AB Mitogen-activated protein (MAP) kinaseactivated protein kinase 2 (MK2) is one of

several kinases directly regulated by p38 MAP kinase. A role of p38 MAP kinase in ischemic brain injury has been previously suggested by pharmacological means. In the present study, we provide evidence for a role of MK2 in cerebral ischemic injury using MK2-deficient (MK2(-/-)) mice. MK2(-/-) mice subjected to focal ischemia markedly reduced infarct size by 64 and 76% after transient and permanent ischemia, respectively, compared with wild-type mice. Furthermore, MK2(-/-) mice had significant reduction in neurological deficits. Real-time PCR analysis identified a significantly lower expression in interleukin-1beta mRNA (53% reduction) but not in tumor necrosis factor-alpha mRNA in MK2(-/-) mice over wild-type animals after ischemic injury. The significant reduction in interleukin-1beta was also confirmed in MK2(-/-) mice by enzyme-linked immunosorbent assay. The marked neuroprotection from ischemic brain injury in MK2(-/-) mice was not associated with the alteration of hemodynamic or systemic variables, activation of caspase-3, or apoptosis. Our data provide new evidence for the involvement of MAP kinase pathway in focal ischemic brain injury and suggest that this effect might be associated with the expression of interleukin-1beta in the ischemic brain tissue.

- L10 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
- AB Myocardial adaptation to **ischemia** has been shown to activate protein tyrosine kinase, potentiating activation of phospholipase D, which leads to the stimulation of mitogen-activated protein (MAP) kinases and MAP kinase-activated protein (MAPKAP) kinase 2. The present study sought

to further examine the signal transduction pathway for the MAPKAP kinase 2 activation during ischemic adaptation. Isolated perfused rat hearts were adapted to ischemic stress by repeated ischemia and reperfusion. Hearts were pretreated with genistein to block tyrosine kinase, whereas SB-203580 was used to inhibit p38 MAP kinases. Western blot anal. demonstrated that p38 MAP kinase is phosphorylated during ischemic stress adaptation. Phosphorylation of p38 MAP kinase was blocked by genistein, suggesting that activation of p38 MAP kinase during ischemic adaptation is mediated by a tyrosine kinase signaling pathway. MAPKAP kinase 2 was estimated by following in vitro phosphorylation with recombinant human heat shock protein 27 as specific substrate for MAPKAP kinase 2. Again, both genistein and SB-203580 blocked the activation of MAPKAP kinase 2 during myocardial adaptation to ischemia. Immunofluorescence microscopy with anti-p38-antibody revealed that p38 MAP kinase is primarily localized in perinuclear regions. P38 MAP kinase moves to the nucleus after ischemic stress adaptation. After ischemia and reperfusion, cytoplasmic striations in the myocytes become obvious, indicating translocation of p38 MAP kinase from nucleus to cytoplasm. Corroborating these results, myocardial adaptation to ischemia improved the left ventricular functions and reduced myocardial infarction that were reversed by blocking either tyrosine kinase or p38 MAP kinase. These results demonstrate that myocardial adaptation to ischemia triggers a tyrosine kinase-regulated signaling pathway, leading to the translocation and activation of p38 MAP kinase and implicating a role for MAPKAP kinase 2.

L10 ANSWER 8 OF 8 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on STN

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L10 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN
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AN 2005:71066 CAPLUS

DN 142:170050

TI DEF domain-containing members of the MAP kinase pathway and their use in screening for drug inhibitors

IN Blenis, John; Murphy, Leon O.

PA President and Fellows of Harvard College, USA

SO PCT Int. Appl., 104 pp.

CODEN: PIXXD2

DT Patent

LA English

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	PATENT NO.					KIND		DATE		APPLICATION NO.						DATE			
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L10 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

DN 140:229441

AN 2004:182659 CAPLUS

TI Methods of reducing ischemic injury with compounds reducing activity of MAP kinase-activated protein

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IN
     Wang, Xinkang; Schieven, Gary; Feuerstein, Giora Z.
PA
     Bristol-Myers Squibb Company, USA
     PCT Int. Appl., 39 pp.
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     PubMed ID: 12215446
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     Mitogen-activated protein kinase-activated protein (MAPKAP) kinase 2
     deficiency protects brain from ischemic injury in mice.
ΑU
     Wang Xinkang; Xu Lin; Wang Hugh; Young Peter R; Gaestel Matthias;
     Feuerstein Giora Z
     Department of Cardiovascular Sciences, Bristol-Myers Squibb Company,
CS
     Wilmington, Delaware 19880-0400, USA.. xinkang.wang@bms.com
SO
     The Journal of biological chemistry, (2002 Nov 15) Vol. 277, No. 46, pp.
     43968-72. Electronic Publication: 2002-09-04.
     Journal code: 2985121R. ISSN: 0021-9258.
CY
     United States
DT
     Journal; Article; (JOURNAL ARTICLE)
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FS
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     200301
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     Entered STN: 13 Nov 2002
     Last Updated on STN: 3 Jan 2003
     Entered Medline: 2 Jan 2003
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     1998:759299 CAPLUS
DN
     130:123195
TI
     Ischemic preconditioning triggers tyrosine kinase signaling: a potential
     role for MAPKAP kinase 2
ΑU
     Maulik, Nilanjana; Yoshida, Tetsuya; Zu, You-Li; Sato, Motoaki; Banerjee,
     Anirban; Das, Dipak K.
CS
     Departments of Surgery and Physiology, University of Connecticut School of
     Medicine, Farmington, CT, 06030-1110, USA
SO
     American Journal of Physiology (1998), 275(5, Pt. 2), H1857-H1864
     CODEN: AJPHAP; ISSN: 0002-9513
PB
     American Physiological Society
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kinase 2

DT Journal

LA English

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AN 1997:424888 BIOSIS

DN PREV199799724091

TI Involvement of a tyrosine kinase-dependent signal transduction process involving p-38 MAP kinases and MAPKAP kinase 2 in ischemic preconditioning.

AU Das, Dipak K.

CS Univ. Connecticut Sch. Med., Farmington, CT, USA

SO Journal of Molecular and Cellular Cardiology, (1997) Vol. 29, No. 7, pp. A272.

Meeting Info.: XIX Annual Meeting of the International Society for Heart Research (American Section) Cardiovascular Injury, Repair and Adaptation. Vancouver, British Columbia, Canada. July 23-27, 1997.

CODEN: JMCDAY. ISSN: 0022-2828.

DT Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LA English

ED Entered STN: 8 Oct 1997

Last Updated on STN: 21 Nov 1997

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